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                 PCTGEN now available on STN
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         Feb 24
                 TEMA now available on STN
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         Feb 26 PCTFULL now contains images
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                 structures available in REGISTRY
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                 MEDLINE Reload
NEWS 12
         Apr 17
                 Polymer searching in REGISTRY enhanced
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                 Indexing from 1927 to 1936 added to records in CA/CAPLUS
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         Apr 21
                 New current-awareness alert (SDI) frequency in
                 WPIDS/WPINDEX/WPIX
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         Apr 28
                 RDISCLOSURE now available on STN
NEWS 16
         May 05
                 Pharmacokinetic information and systematic chemical names
                 added to PHAR
NEWS 17
         May 15
                 MEDLINE file segment of TOXCENTER reloaded
NEWS 18
         May 15
                 Supporter information for ENCOMPPAT and ENCOMPLIT updated
NEWS 19
         May 19
                 Simultaneous left and right truncation added to WSCA
NEWS 20
         May 19
                 RAPRA enhanced with new search field, simultaneous left and
                 right truncation
NEWS 21
         Jun 06
                 Simultaneous left and right truncation added to CBNB
NEWS 22
         Jun 06
                 PASCAL enhanced with additional data
NEWS 23
         Jun 20
                 2003 edition of the FSTA Thesaurus is now available
NEWS 24
         Jun 25
                 HSDB has been reloaded
NEWS 25
         Jul 16
                 Data from 1960-1976 added to RDISCLOSURE
NEWS 26
         Jul 21
                 Identification of STN records implemented
NEWS 27
         Jul 21
                 Polymer class term count added to REGISTRY
NEWS 28
                 INPADOC: Basic index (/BI) enhanced; Simultaneous Left and
         Jul 22
                 Right Truncation available
         AUG 05
NEWS 29
                 New pricing for EUROPATFULL and PCTFULL effective
                 August 1, 2003
NEWS 30
         AUG 13
                 Field Availability (/FA) field enhanced in BEILSTEIN
NEWS 31
         AUG 15
                 PATDPAFULL: one FREE connect hour, per account, in
                 September 2003
NEWS 32
         AUG 15
                 PCTGEN: one FREE connect hour, per account, in
                 September 2003
NEWS 33
         AUG 15
                 RDISCLOSURE: one FREE connect hour, per account, in
                 September 2003
NEWS 34
         AUG 15
                 TEMA: one FREE connect hour, per account, in
                 September 2003
                 Data available for download as a PDF in RDISCLOSURE
NEWS 35
         AUG 18
NEWS 36
         AUG 18
                 Simultaneous left and right truncation added to PASCAL
NEWS 37
         AUG 18
                 FROSTI and KOSMET enhanced with Simultaneous Left and Right
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NEWS 38 AUG 18 Simultaneous left and right truncation added to ANABSTR

NEWS EXPRESS April 4 CURRENT WINDOWS VERSION IS V6.01a, CURRENT MACINTOSH VERSION IS V6.0b(ENG) AND V6.0Jb(JP),

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This file contains CAS Registry Numbers for easy and accurate substance identification.

=> pronase and virus (w) isolation

7654 PRONASE

13 PRONASES

7661 PRONASE

(PRONASE OR PRONASES)

286631 VIRUS

61896 VIRUSES

296997 VIRUS

(VIRUS OR VIRUSES)

219875 ISOLATION

947 ISOLATIONS

220482 ISOLATION

```
(ISOLATION OR ISOLATIONS)
          1021 VIRUS (W) ISOLATION
             2 PRONASE AND VIRUS (W) ISOLATION
L1
=> (streptomyces griseus trypsin)
         32009 STREPTOMYCES
          4221 GRISEUS
         64422 TRYPSIN
           473 TRYPSINS
         64464 TRYPSIN
                  (TRYPSIN OR TRYPSINS)
L2
            61 (STREPTOMYCES GRISEUS TRYPSIN)
                  (STREPTOMYCES (W) GRISEUS (W) TRYPSIN)
=> virus (w) putification
        286631 VIRUS
         61896 VIRUSES
        296997 VIRUS
                  (VIRUS OR VIRUSES)
             O PUTIFICATION
L3
             O VIRUS (W) PUTIFICATION
=> Virus (w) purification
        286631 VIRUS
         61896 VIRUSES
        296997 VIRUS
                 (VIRUS OR VIRUSES)
        289567 PURIFICATION
           874 PURIFICATIONS
        290151 PURIFICATION
                  (PURIFICATION OR PURIFICATIONS)
        249592 PURIFN
           232 PURIFNS
        249695 PURIFN
                  (PURIFN OR PURIFNS)
        419722 PURIFICATION
                  (PURIFICATION OR PURIFN)
L4
           915 VIRUS (W) PURIFICATION
=> L4 and L2
L5
             0 L4 AND L2
=> virus (w) isolation
        286631 VIRUS
         61896 VIRUSES
        296997 VIRUS
                  (VIRUS OR VIRUSES)
        219875 ISOLATION
           947 ISOLATIONS
        220482 ISOLATION
                 (ISOLATION OR ISOLATIONS)
L6
          1021 VIRUS (W) ISOLATION
=> L6 and L2
L7
             0 L6 AND L2
=> pronase
          7654 PRONASE
            13 PRONASES
          7661 PRONASE
rs
                  (PRONASE OR PRONASES)
```

=> L6 and L8

L9

2 L6 AND L8

=> L8 and L4

L10 8 L8 AND L4

=> L10 and HAV

1026 HAV

19 HAVS

1033 HAV

(HAV OR HAVS)

L11

0 L10 AND HAV

=> DIS L10 1- IBIB ABS

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L10 ANSWER 1 OF 8 CAPLUS COPYRIGHT 2003 ACS on STN

ACCESSION NUMBER:

1997:270740 CAPLUS

DOCUMENT NUMBER:

126:248758

TITLE:

Purification and crystallization of the attachment

proteins of enveloped animal viruses using a virosome

intermediate

INVENTOR(S):

Portner, Allen; Takimoto, Toru

PATENT ASSIGNEE(S):

St. Jude Children's Research Hospital, USA; Portner,

Allen; Takimoto, Toru PCT Int. Appl., 40 pp.

SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

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PATENT NO.
                  KIND DATE
                                      APPLICATION NO. DATE
                   A1 19970313 WO 1996-US14187 19960906
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    WO 9709345
        W: AL, AM, AT, AU, AZ, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK,
           EE, ES, FI, GB, GE, HU, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, .
           LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO,
           RU, SD, SE, SG, SI, SK, TJ, TM, TR, TT, UA, UG, US, UZ, VN, AM,
           AZ, BY, KG, KZ, MD, RU, TJ, TM
        RW: KE, LS, MW, SD, SZ, UG, AT, BE, CH, DE, DK, ES, FI, FR, GB, GR,
            IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI
    AU 9671545
                    A1 19970327
                                       AU 1996-71545
                                                       19960906
PRIORITY APPLN. INFO.:
                                    US 1995-3447P P 19950908
                                    WO 1996-US14187 W 19960906
```

AB A method of purifying the attachment proteins of enveloped viruses in a biol. active form suitable for crystn. and X-ray crystallog. anal. is described. The proteins are incorporated into virosomes by solubilization of the virus with detergent followed by sedimentation of the nucleocapsid and matrix proteins. Th supernatant contg. the solubilized attachment proteins and envelope lipids is then treated to remove the detergent with reconstitution of virosomes. The sol. domains of the protein can then be removed by proteolytic cleavage and the residual virosomes removed by sedimentation. The hemagglutinin-neuraminidase (HN) of the Kansas strain of Newcastle's disease virus was purified from allantoic fluid by resuspending the virus at 20 mg/mL in PBS contg. Triton X-100 2 vol.% and incubated at room temp. for 1 h. Nucleocapsid and matrix proteins were pelleted by centrifugation and detergent removed from the supernatant using Bio-Beads. The extracellular domain of the HN was solubilized by treatment with pronase to give a single band on gel electrophoresis.

L10 ANSWER 2 OF 8 CAPLUS COPYRIGHT 2003 ACS on STN ACCESSION NUMBER: 1991:467248 CAPLUS

DOCUMENT NUMBER:

115:67248

TITLE:

A new method for the purification of the influenza A

virus neuraminidase

AUTHOR (S):

McKimm-Breschkin, J. L.; Caldwell, J. B.; Guthrie, R.

E.; Kortt, A. A.

CORPORATE SOURCE:

Div. Biomol. Eng., CSIRO, Parkville, 3052, Australia Journal of Virological Methods (1991), 32(1), 121-4

CODEN: JVMEDH; ISSN: 0166-0934

DOCUMENT TYPE:

Journal

LANGUAGE:

SOURCE:

English

A rapid new method for the purifn. of neuraminidase (NA) heads from influenza A virus is described. Virus was pelleted directly from allantoic fluid and was digested with Pronase. The cores were removed by centrifugation, redigested and the released NA heads were pooled and concd. The NA was sepd. from all contaminating proteins in a single step on a Superose 12 column. The purified material was suitable for both crystallog. and for the prodn. of monospecific antisera.

L10 ANSWER 3 OF 8 CAPLUS COPYRIGHT 2003 ACS on STN

ACCESSION NUMBER:

1984:402572 CAPLUS

DOCUMENT NUMBER:

101:2572

TITLE:

Partial characterization of a transformation-specific

glycopeptide in SSV-NP cells

AUTHOR (S):

Thiel, Heinz Juergen; Hafenrichter, Rudolf; Greger,

Bernd

CORPORATE SOURCE:

Fed. Res. Cent. Virus Dis. Anim., Tuebingen, D-7400,

Fed. Rep. Ger.

SOURCE:

AB

Virology (1984), 134(1), 138-47 CODEN: VIRLAX; ISSN: 0042-6822

DOCUMENT TYPE:

Journal English

LANGUAGE:

An autologous antiserum against simian sarcoma virus-infected nonproducer cells (SSV-NP cells) recognized a SSV transformation-specific glycopeptide (SSV-TrSgp) (Thiel, H. J., et al., 1981). Gel filtration of this component on a Sephacryl S-200 column indicated an apparent mol. wt. at .apprx.200,000. This antigen represented a proteoglycan-like mol., as evidenced by the size of glycopeptides after Pronase treatment and by incubation with chondroitinases. The antigenicity of the SSV-TrSgp was completely destroyed after exposure to different proteases. On the other hand, incubation with neuraminidase or chondroitinases degraded the mol. to some extent, but did not affect its antigenicity as measured by immunopptn. Trypsin and EDTA treatment of intact pulse-labeled cells, as well as surface iodination, indicated that the SSV-TrSqp represents a cell membrane-assocd. mol.

L10 ANSWER 4 OF 8 CAPLUS COPYRIGHT 2003 ACS on STN

1980 123603 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER:

92:123603

TITLE:

Isolation and preliminary characterization of herpes

Channel Catfish virus DNA

Robin, Jean; Rodrigue, Alice

CORPORATE SOURCE: Fac. Sci., Univ. Sherbrooke, Sherbrooke, QC, J1K 2R1,

SOURCE:

Canadian-Journal of Microbiology (1980), 26(2), 130-4

CODEN: CJMIAZ; ISSN: 0008-4166

DOCUMENT TYPE:

Journal

LANGUAGE:

AUTHOR(S):

English

The DNA of Channel Catfish virus (CCV) was selectively extd. from infected cells with a 5% soln. of Na deoxycholate, deproteinized with Na sarcosinate and Pronase, and purified by PhOH extn. followed by equil. d. gradient centrifugation in a CsCl soln. CCV DNA displayed a buoyant d. of 1.715 g/cm3 in such a soln., as would be expected from a duplex DNA contg. 56.1% guanine plus cytosine. As estd. from both its sedimentation coeff. and length in the electron microscope, CCV DNA is a

linear duplex mol. of .apprx.85 .times. 106 daltons.

L10 ANSWER 5 OF 8 CAPLUS COPYRIGHT 2003 ACS on STN

ACCESSION NUMBER: 1978:147999 CAPLUS

DOCUMENT NUMBER: 88:147999

TITLE: Isolation and study of the electrophoretic mobility of

pig influenza virus neuraminidase

AUTHOR(S): Tolmacheva, V. P.; Daulbaeva, K. D.; Isaeva, E. S.;

Chuvakova, Z. K.; Amantaev, S. Zh.

CORPORATE SOURCE: USSR

SOURCE: Izvestiya Akademii Nauk Kazakhskoi SSR, Seriya

Biologicheskaya (1978), 16(1), 51-5

CODEN: IKABAR; ISSN: 0002-3183

DOCUMENT TYPE: Journal LANGUAGE: Russian

AB Neuraminidase from pig influenza virus was isolated by treatment of virus with ether, Tween 20, and pronase followed by centrifugation at 45,000 rpm for 2 h. The products of virus disintegration were pptd. by treatment with formalinized erythrocytes and neuraminidase was found in the supernatant liq. The decrease in yield after erythrocyte treatment indicates that a significant portion of the enzyme is found in complexes with hemagglutinins, which are easily adsorbed on erythrocytes. Polyacrylamide gel electrophoresis showed 2 components with neuraminidase activity; the 1st component decreased significantly after treatment with erythrocytes. The mol. wts. were 250,000 and 200,000 daltons, characteristic for a tetrameric structure of neuraminidase.

L10 ANSWER 6 OF 8 CAPLUS COPYRIGHT 2003 ACS on STN

ACCESSION NUMBER: 1977:417649 CAPLUS

DOCUMENT NUMBER: 87:17649

TITLE: Partial purification and characterization of the

potato virus Y helper component

AUTHOR(S): Govier, D. A.; Kassanis, B.; Pirone, T. P. CORPORATE SOURCE: Rothamsted Exp. Stn., Harpenden/Herts., UK

SOURCE: Virology (1977), 78(1), 306-14 CODEN: VIRLAX; ISSN: 0042-6822

DOCUMENT TYPE: Journal LANGUAGE: English

AB Mg2+ stabilized potato virus Y helper component during partial purifn. In solns. contg. 0.02M Mg2+, the helper component retained most of its activity for 2 days at 4.degree. and for .gtoreg.8 months at -15.degree.. Activity was destroyed on incubation with Pronase-or-trypsin or by heating for 5 min at 55.degree., but not by incubation with RNase. Incubation with its own antiserum strongly inhibited helper component activity, but antisera to potato virus Y coat protein or inclusion protein had no more effect than a control serum. Filtration through a Sephadex G-200 column resulted in a broad peak of activity which produced many protein-staining bands when electrophoresed on polyacrylamide gel. Gel filtration and ultrafiltration expts. both indicated a mol. wt. of 100,000-200,000. Some helper component activity was retained by aphids allowed to probe into a sucrose soln. for 20 min, showing that the helper component is more firmly bound to the aphid than is the tobacco mosaic virus-poly-L-ornithine complex.

L10 ANSWER 7 OF 8 CAPLUS COPYRIGHT 2003 ACS on STN

ACCESSION NUMBER: 1977:402099 CAPLUS

DOCUMENT NUMBER: 87:2099

TITLE: Highly infectious RNA isolated from cowpea chlorotic

mottle virus with low specific infectivity

AUTHOR(S): Wyatt, S. D.; Kuhn, C. W.

CORPORATE SOURCE: Dep. Plant Pathol. Plant Genet., Univ. Georgia,

Athens, GA, USA

SOURCE: Journal of General Virology (1977), 35, Pt. 1, 175-80

CODEN:_JGVIAY;_ISSN:_0022=1317

DOCUMENT TYPE: Journal LANGUAGE: English

AB Recovery and specific infectivity of infectious RNA from cowpea chlorotic mottle virus of low specific infectivity (14-21 day infections) were greatly improved by using antioxidants during virus purifn. and RNA extn., and by disrupting coat protein with pronase before PhOH-Na dodecyl sulfate extn. Total infectivity of RNA from virus of low infectivity was increased >30-fold. RNA profiles obtained using polyacrylamide gels were then similar for virus with high (4-7 day infections) or low specific infectivity. Low specific invectivity, therefore, seems to be caused by alteration of the coat protein or of the protein-RNA interaction in intact virus particles.

L10 ANSWER 8 OF 8 CAPLUS COPYRIGHT 2003 ACS on STN

ACCESSION NUMBER: 1974:105218 CAPLUS

DOCUMENT NUMBER: 80:105218

TITLE: Preparative isolation of neuraminidase from influenza

A viruses (Singapore) 1/57, (Hong Kong) 1/68, and

(Leningrad) 99/71

AUTHOR(S): Simanovskaya, V. K.; Vaitkiene, V.; Golubev, D. B.

CORPORATE SOURCE: Vses. Nauchno-Issled. Inst. Grippa, Leningrad, USSR

SOURCE: Voprosy Virusologii (1973), (5), 555-60

CODEN: VVIRAT; ISSN: 0507-4088

DOCUMENT TYPE: Journal LANGUAGE: Russian

AB The 3 title viruses cultured in the allantoic fluid of chick embryos were used as sources for the prepn. of neuraminidase (I). The extn. method comprised disintegration of the virus with BuOH, ether, and Pronase, pptn. of the S antigen, and gel filtration on Sephadex G-200. The purified I was homogeneous upon polyacrylamide gel electrophoresis. The Singapore strain gave a mugh higher yield of I, with a much higher sp. activity than either of the other 2 strains. In the process of purification, I acquired a gradually increasing specificity toward the low-mol.-wt. compd. sialyllactose, as opposed to the high-mol.-wt. ovomucin.

=> DIS L9 1- IBIB ABS

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L9 ANSWER 1 OF 2 CAPLUS COPYRIGHT 2003 ACS on STN

ACCESSION NUMBER: 1985:75385 CAPLUS

DOCUMENT NUMBER: 102:75385

TITLE: Pronase treatment of type A/H2N2/ and type B

influenza viruses. Isolation of

pure neuraminidase heads

AUTHOR(S): Kavaklova, L.; Praskov, D.; Vulkova, B.; Petrunova,

S.; Nikolova, Z.; Kotseva, R.

CORPORATE SOURCE: Med. Akad., Sofia, Bulg.

SOURCE: Epidemiologiya, Mikrobiologiya i Infektsiozni Bolesti

(1984), 21(4), 23-31

CODEN: EMIBA3; ISSN: 0425-1482

DOCUMENT TYPE: Journal

LANGUAGE: (Bulgarian)

AB The direct effect of pronase, a protease, was studied on 4 type A and 1 type B influenza virus stains. Pure and active neuramindase with a very good yield was isolated from strain A/Singapore/1/57/H2N2/. The other 3 type A/H3N2/ strains appeared to have thermolabile and pronase-sensitive neuraminidase and a hemagglutinin relatively resistant to pronase degrdn. The neuraminidase prepns. isolated had low enzyme activity and were hemagglutinin-polluted. Pure neuraminidase with reduced enzyme activity was isolated from strain

B/Singapore 222/79. Monospecific antisera against the pure neuraminidase heads, isolated from A/Singapore/1/57 and B/Singapore 222/79, were obtained. The antisera were used in the double agar diffusion test, aimed at comparing the antigenic identity of type N2 and type B neuraminidases accordingly.

ANSWER 2 OF 2 CAPLUS COPYRIGHT 2003 ACS on STN

ACCESSION NUMBER: 1982:612236 CAPLUS

DOCUMENT NUMBER: 97:212236

TITLE: Isolation of native alpha-virus RNA and several of its

physicochemical indexes

Uryvaev, L. V.; Klimenko, S. M.; Samokhvalov, E. I.; AUTHOR (S):

Iferov, V. P.

CORPORATE SOURCE: Inst. Virusol. im. Ivanovskogo, Moscow, USSR

SOURCE: Deposited Doc. (1981), VINITI 4502-81, 18 pp. Avail.:

> VINITI Report

DOCUMENT TYPE: LANGUAGE: Russian

Genomic-RNA was isolated from Venezuelan equine encephalomyelitis virus, propagated in chick embryo fibroblasts, by extn. with phenol, treatment with detergent (SDS), or promaser the atment. The yield of RNA was 80-95%. Anal. of RNA in sucrose d. gradient and gel electrophoresis revealed the presence of 3 RNA species with sedimentation coeffs. of 42 S, 28 S, and 18 S. The viral RNA had a mol. wt. of 4.0-4.1 megadaltons and was composed of 11,500-12,500 nucleotides. Electron micrographs of the viral genome are given.